Wintertime aerosol optical properties in Lanzhou, Northwest China: Emphasis on the rapid increase of aerosol absorption under high particulate pollution

Xu Guan, Min Wang, Tao Du, Pengfei Tian, Naiyue Zhang, Jinsen Shi, Yi Chang, Lei Zhang, Min Zhang, Xin Song, Yijia Sun

Key Laboratory for Semi-Arid Climate Change of the Ministry of Education, College of Atmospheric Sciences, Lanzhou University, Lanzhou, 730000, China
Collaborative Innovation Center for Western Ecological Safety, Lanzhou University, Lanzhou, 730000, China
Gansu Province Environmental Monitoring Center, Lanzhou, 730020, China

HIGHLIGHTS
- Winter aerosol absorption increased under high particulate pollution in Lanzhou.
- Elemental carbon and fine soils accounted for most of the aerosol absorption.
- Radiative forcing efficiency was highly correlated with single scattering albedo.

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ABSTRACT
The evolution of aerosol absorption and the contribution of absorbing species under different severities of particulate pollution are poorly understood, though absorption is key in aerosol radiative forcing. To resolve the problems, aerosol absorbing properties from low to high particulate pollution were investigated by using intensive observations of aerosol optical properties in the winter of 2019–2020 in Lanzhou, Northwest China. The aerosol scattering coefficient increased linearly with increasing particulate matter <2.5 μm in diameter (PM$_{2.5}$) and the absorption coefficient increased more rapidly under higher particulate pollution, leading to rapid decline in single scattering albedo (SSA) and sharp increase in mass absorption efficiency of PM$_{2.5}$ (MAE$_{PM2.5}$). The SSA (MAE$_{PM2.5}$) decreased (increased) from 0.87 (0.76) in the lowest PM$_{2.5}$ bin to 0.82 (1.11) in the highest PM$_{2.5}$ bin. The linear relationship between the scattering coefficient and PM$_{2.5}$ was attributed to decreasing aerosol hygroscopicity with increasing PM$_{2.5}$. Elemental carbon (EC), fine soils (FS), and organic carbon (OC) accounted for 77.4%, 16.6%, and 6.0% of the total aerosol absorption, respectively. From low to high particulate pollution levels, the contribution of EC absorption increased from 68.3% to 80.5% while that of FS decreased from 25.5% to 13.9%. The aerosol radiative forcing efficiency was strongly correlated with SSA. Our results show a unique rapid increase in aerosol absorption under high particulate pollution during winter in Lanzhou, which is opposite to the trends observed in eastern Chinese cities, where SSA increases with increasing PM$_{2.5}$.

1. Introduction

Atmospheric aerosols impact the radiative budget of Earth’s atmospheric system via aerosol–radiation and aerosol–cloud interactions (Twomey, 1977; Haywood and Ramaswamy, 1998; Bellouin et al., 2020). Absorbing aerosols play a unique role in aerosol–radiation interactions due to the heating effect in and at the top of the atmosphere (Jacobson, 2001). Specifically, absorbing aerosols deteriorate air quality (Ding et al., 2016; Zhang et al., 2020a), increase extreme weather (Fan et al., 2015), impact global and regional climate change (Bond et al., 2013), and damage human health (Janssen et al., 2011). Absorbing aerosols are increasingly of interest to researchers and policymakers (IPCC, 2013; Liu et al., 2020). Relevant emission reduction policies have been introduced for absorbing aerosols such as black carbon (Shindell...
et al., 2012). However, assessing the radiative effects of absorbing aerosols remains challenging due to various reasons such as uncertainty of emission sources (Ferrero et al., 2018), complex optical properties (Andreae and Gelencser, 2006; Tian et al., 2018), methodological consistency and sophistication in observations (Yu et al., 2006; Barman et al., 2019), spatial and temporal variations (Rang et al., 2017), and poor parameterization in model simulations (Liu et al., 2017).

Absorbing aerosols heat the atmosphere and generally have a positive radiative force on the climate system. The global aerosol radiative absorption has been estimated to vary from 0.1 to 0.6 W m\(^{-2}\) (Allen et al., 2012; Myhre et al., 2013) to 0.9–1.2 W m\(^{-2}\) (Sato et al., 2003; Ramanathan and Carmichael, 2008). Black carbon (BC) is the major absorbing aerosol species (Bond et al., 2006; Cappa et al., 2012) and BC in polluted urban environments plays an essential role in pollution development and greatly contributes to large positive radiative forcing (Feng et al., 2016). Mineral dust and part of organic carbon (OC) are also absorbing aerosols in addition to BC. Mineral dust affects the radiation balance of Earth’s atmospheric system via interactions with solar short-wave radiation and Earth’s long-wave radiation (Miller and Tegen, 1998; Hu et al., 2019). It has been estimated that more than half of the aerosol absorption optical depth at 550 nm is derived from dust aerosols (Chin et al., 2009). Absorbing OC (or brown carbon, BrC) mainly originates from the combustion of coal and biomass and exhibits significant absorption at the ultraviolet or shorter wavelengths (Feng et al., 2013; Bergin et al., 2015). The contribution of BC to the absorption aerosol optical depth for carbonaceous aerosols is greater in urban sites than in the background areas, while the contribution of BrC is higher in background sites (Cho et al., 2019). It was found that the contribution of BrC to the total light absorption at 635 nm can be as much as 34% in Lanzhou (Zhang et al., 2019). However, our understanding of the contributions of BC, mineral dust, and OC to absorption under different severities of particulate pollution remains limited.

The fraction of aerosol species varies under different severities of particulate pollution, which leads to changing absorption in complex urban aerosols. The absorbing aerosols (i.e., BC, dust, and BrC) are mainly derived from primary emissions (Bond et al., 2006; Bergin et al., 2015; Hu et al., 2019), while secondarily generated aerosols (e.g., sulfate, nitrate, and ammonium) contribute greatly to light scattering (Liu et al., 2008; Tao et al., 2015). The aging of BC aerosols causes absorption enhancement due to the “lensing effect” (Jacobson, 2001). High relative humidity promotes aerosol hygroscopicity growth and aqueous chemical reactions (Svenningsson et al., 1992; Wang et al., 2016), which is favorable for light scattering enhancement (Titos et al., 2016; Zhao et al., 2019). Although the absorption of aerosols in East China has increased since 2013 (Tao et al., 2020), the variation in the aerosol absorption for different particulate pollution is still unknown. For example, the single scattering albedo (SSA) in Beijing decreased moderately from 0.82 in 2014 to 0.78 in 2016 (Wang et al., 2019) and also showed an increasing trend during pollution events (Yang et al., 2015). The average SSA was 0.91 under clean conditions and increased to 0.93 during polluted days in Wuhan (Liao et al., 2020). However, observations from Guangdong revealed that the SSA decreased from 0.93 to 0.91 with increasing particulate pollution (Kong et al., 2020). Single scattering albedo also varies among cities and seasons (Cao et al., 2012; Segura et al., 2016; Venter et al., 2020). Overall, the evolution of aerosol absorption under different severities of particulate pollution has been poorly quantified and the reasons for the existing discrepancies in aerosol absorption among the cities worldwide remain unclear.

The fraction of secondary inorganic aerosol has been observed to decrease with increasing particulate matter <2.5 μm in diameter (PM\(_{2.5}\)) in Lanzhou (Du et al., 2020), which is opposite to results in eastern Chinese cities, such as Beijing, where aerosol scattering coefficient exhibits an exponential growth with PM\(_{2.5}\) due to high hygroscopicity and strong aqueous chemical reactions (Zhang et al., 2020b). Lanzhou is in a semi-arid region that is characterized by low relative humidity and weak aqueous chemical reactions during winter (Du et al., 2020), which is unfavorable for light scattering enhancement. Furthermore, emissions of absorbing aerosols, including elemental carbon (EC), mineral dust, and OC greatly contribute to aerosol absorption during winter. Thus, the evolution of aerosol absorption under different severities of particulate pollution and the contributions of absorbing species were investigated based on multiple observations of aerosol optical properties in the winter of 2019–2020 in Lanzhou. Firstly, the aerosol optical properties were overviewed and found that SSA is relatively low in Lanzhou when compared with eastern Chinese cities. Then the features of aerosol scattering and absorption from low to high particulate pollution and the contribution of aerosol species were studied and results showed that absorption under high particulate pollution increased rapidly. Finally, the influence of aerosol species on aerosol absorption during wintertime in Lanzhou was analyzed.

2. Data and methodology

2.1. Overview of data

Field measurements were performed from December 3, 2019 to February 4, 2020 at the Lanzhou Atmospheric Components Monitoring Superstation (LACMS; 36.05°N, 103.87°E, 1520 m a.s.l.), which is on the campus of Lanzhou University at downtown area of Lanzhou City. Multiple online instruments were deployed to continuously measure the concentrations of particulate matter (PM) of different sizes, aerosol optical properties, chemical compositions, gases, atmospheric visibility, and meteorological parameters. An overview of the instruments is provided in Table S1. Detailed introduction of the site is available elsewhere (Du et al., 2020).

A PM synchronous mixing monitor (5030i; SHARP, Thermo Fisher, USA) was used to monitor the hourly mass concentrations of PM with an aerodynamic diameter of up to 1.0 μm (PM\(_{1.0}\)), 2.5 μm (PM\(_{2.5}\)), 10 μm (PM\(_{10}\)), and total suspended particles (TSP) with a flow rate of 1 m\(^3\) h\(^{-1}\). Aerosol scattering coefficients of PM\(_{2.5}\) at 450, 520, and 700 nm wavelengths were observed by using a nephelometer (Aurora-3000, BMET Co., Ltd., China). An external heater was placed at the air sampling inlet of the nephelometer, which maintained the temperature of air samples between 50 °C and 60 °C. The nephelometer also contained an internal heater, which heated the incoming air samples when their relative humidity (RH) of the air samples exceeded 40%. The two heaters evaporated the water droplets and decreased the RH to <20% at the LACMS site, which was measured by an integrated humidity sensor and displayed on the main screen of the nephelometer.

An aethalometer (AE33, Magee Scientific, USA) with a PM\(_{2.5}\) inlet was used to measure the aerosol light absorption coefficients at 370, 470, 520, 590, 660, 880, and 950 nm. The matrix scattering effect of the AE33 aethalometer filter was corrected using the matrix scattering effect correction coefficient, C = 1.39, provided in the instruction manual for the filter tape. The influence of RH on the absorption coefficient is discussed in the Supplementary Materials (Text S2 and Fig. S5). No relationship was found between the absorption coefficient and RH.

The hourly water-soluble inorganic ions (Cl\(^–\), NO\(_3\)\(^–\), SO\(_4\)\(^{2–}\), NH\(_4\)\(^+\), Na\(^+\), K\(^+\), Mg\(^2+\), and Ca\(^{2+}\)) in PM\(_{2.5}\) were measured using a Monitor for Aerosols and Gases in Ambient Air (MARGA, Applikon Biotechnology B. V., The Netherlands), an advanced and widely used instrument worldwide (Brink et al., 2019). An online and continuous particulate carbon monitor (Model-4/OC/EC (RT-4) Lab, Sunset Laboratory, Inc., USA) was used to measure the hourly OC and EC in PM\(_{2.5}\). This OC/EC monitor adopts the thermal/optical method approved by National Institute for Occupational Safety and Health to measure OC and EC collected on a quartz filter membrane.

Concentrations of trace gases, including O\(_3\), SO\(_2\), NO, NO\(_2\), and CO, were obtained at a resolution of 1 h using a set of online gas analyzers (491, 43i, 42i, and 48i, respectively; Thermo Fisher Scientific, USA). Hourly meteorological parameters were collected using a meteorological sensor (FWS500, Fronttech, Ltd., China), including temperature,
RH, air pressure, wind direction and wind speed. Atmospheric visibility at 880 nm was measured using a visibility meter (DNQ-2, China).

To ensure the quality of the observation data, all instruments were installed and operated in strict accordance with their instruction manuals and were calibrated according to regulations on time. Details on the calibrations are available in Table S3.

To better understand the changes in the optical properties and chemical compositions of aerosols under different severities of particulate pollution, the observations were grouped using two methods. First, observations were grouped into low (daily PM$_{2.5}$ ≤ 35 μg m$^{-3}$), moderate (35 < daily PM$_{2.5}$ ≤ 75 μg m$^{-3}$), and high (daily PM$_{2.5}$ ≥ 75 μg m$^{-3}$) pollution levels. Sample numbers of the low, moderate, and high PM levels accounted for 9.4%, 40.6%, and 50.0%, respectively of the total samples. We also divided the total samples into 10 bins from low to high particulate pollution, with each bin including 149 samples.

2.2. A brief introduction to aerosol optical parameters

Black carbon is the light-absorbing carbon measured by optical instruments (Petzold et al., 2013). Carbonyl carbon aerosols are divided into OC and EC components via chemical thermo-optical (or thermal) methods. In emissions inventories and climate science, BC is usually synonymous with EC (Cheng et al., 2011). In the present study, BC was used to describe optical performance, while EC was related to chemical species.

The aerosol absorption coefficient was calculated as follows:

$$\sigma_{ap} = \|BC\|_\lambda \times MAC_s$$

where $\|BC\|_\lambda$ is the BC concentration and MAC$_s$ is the cross section of mass absorption. Unless otherwise specified, $\sigma_{ap}$ is the scattering coefficient at 520 nm and $\sigma_{sp}$ is the scattering coefficient at 520 nm. The SSA was calculated from $\sigma_{ap}$ and $\sigma_{sp}$ at 520 nm to be:

$$SSA = \frac{\sigma_{ap}}{\sigma_{ap} + \sigma_{sp}}$$

The scattering Ångström exponent (SAE) is a parameter related to aerosol size; it is relatively low for large particles and high for small particles and may be expressed as follows:

$$SAE = \frac{\log (\sigma_{ap}) - \log (\sigma_{sp})}{\log (\lambda_1) - \log (\lambda_2)}$$

where $\lambda_1 = 520$ and $\lambda_2 = 450$ nm.

The aerosol absorption coefficient is expressed as:

$$\sigma_{ap}(\lambda) = k \cdot \lambda^{-AE}$$

where $k$ is a constant, $\sigma_{ap}(\lambda)$ is the absorption coefficient at wavelength $\lambda$, and AAE is absorption Ångström exponent. AAE was calculated via power-law fitting based on the absorption coefficients at 370, 470, 520, 590, 660, 880, and 950 nm wavelengths. The AAE is an indicator of the dominant light absorber in aerosols. The mass absorption efficiency of PM$_{2.5}$ (MAE$_{PM2.5}$) was calculated as the ratio of the absorption coefficient at 520 nm to the mass concentration of PM$_{2.5}$:

$$MAE_{PM2.5} = \frac{\sigma_{ap}}{PM_{2.5}}$$

The mass scattering efficiency of PM$_{2.5}$ (MSE$_{PM2.5}$) was then calculated by replacing $\sigma_{ap}$ at 520 nm in Equation (5) with $\sigma_{sp}$ at 520 nm.

The atmospheric extinction coefficient ($\sigma_{ext}$,RH) at 550 nm (Mm m$^{-1}$) was calculated based on the visibility (Vis) data as follows (Koschmieder, 1924; Yang et al., 2015):

$$\sigma_{ext}(\lambda) = \frac{3.912 \times 10^3}{\text{Vis}} \times \frac{880}{550}$$

The aerosol scattering coefficient at 550 nm ($\sigma_{sp}$,RH) was calculated as follows:

$$\sigma_{sp}(\lambda) = \sigma_{ext}(\lambda) - \sigma_{ap} - \sigma_{sg} - \sigma_{ag}$$

where $\sigma_{ap}$ is the absorption coefficient at 550 nm, which was calculated from the absorption coefficient at 520 nm by using Equation (4). The scattering coefficient ($\sigma_{sp}$) of the gases was assumed to be a constant of 10 Mm m$^{-1}$ (Liu et al., 2008). The absorption coefficient of gases ($\sigma_{ag}$) was calculated using the experimental equation (Hodkinson, 1966):

$$\sigma_{ag} = [NO_2] \times 0.33$$

where [NO$_2$] is the unit of NO$_2$ (ppbv).

The ratio of the aerosol scattering coefficient at a given RH ($\sigma_{sp}$,RH) to that under dry conditions ($\sigma_{sp}$,dry) (Carriero et al., 2003; Liu et al., 2008):

$$f(RH) = \frac{\sigma_{sp}(\lambda)}{\sigma_{sp}(\lambda) - \sigma_{ap} - \sigma_{sg} - \sigma_{ag}}$$

2.3. Reconstruction of light absorption and scattering coefficients

The light extinction can be expressed as the sum of light scattering by particles ($\sigma_{sp}$) and gases ($\sigma_{sg}$), and the absorption of particles ($\sigma_{ap}$) and gases ($\sigma_{ag}$). The revised Interagency Monitoring of Protected Visual Environments (IMPROVE) algorithm (Pitchford et al., 2007) was used to evaluate the effect of chemical species on aerosol light extinction. The algorithm is detailed in the Supplementary Materials (Text S1).

The PM$_{2.5}$ concentrations were reconstructed by the sum of organic matter (OM), (NH$_4$)$_2$SO$_4$, NH$_4$NO$_3$, EC, fine soil (FS), Na$^+$, and Cl$^-$ to assess whether or not these chemical species were representative of the measured PM$_{2.5}$. A strong positive correlation ($R^2 = 0.93$) was found between the reconstructed and measured PM$_{2.5}$ (Fig. 1a), the regression coefficient was 0.82, indicating that the chemical species were representative of the measured PM$_{2.5}$. A strong positive correlation ($R^2 = 0.92$) was also found between the EC concentration from Sunset OC/EC monitor and concentration by the Magee Scientific AE33 monitor, even though EC was lower than BC (Fig. 1b). The underestimation of EC is further discussed in the Supplementary Materials (Text S3). Since the IMPROVE algorithm assumes that both sulfate and nitrate are present in the form of ammonium sulfate and ammonium nitrate, the presence of sulfate and nitrate in Lanzhou was tested (Fig. 1c). The slope of the linear regression was 1.04, which indicated that ammonium was the
predominant forms of sulfate and nitrate.

The relationship between aerosol chemical species and optical parameters was studied to determine the influence of aerosol composition on optical properties. The scattering coefficient was most strongly correlated with OM and ammonium, followed by nitrate and sulfate (Fig. S2). A strong linear correlation \( (R^2 = 0.93) \) was observed between the absorption coefficient and EC (Fig. 2a) indicating that EC was the main species of absorption. The EC mass absorption coefficient at 520 nm calculated from the linear regression was 24.76 m\(^2\) g\(^{-1}\), which was higher than that of 16.09 m\(^2\) g\(^{-1}\) at 520 nm during summer in Beijing (Zhou et al., 2017). The high EC mass absorption coefficient may be due to deviations from different EC measurement protocols (Chow et al., 2004; Zhi et al., 2011) and the contribution of other absorbing chemical species in addition to EC (Yang et al., 2009).

Aerosol absorption and scattering were reconstructed to quantify the contributions of chemical species via the IMPROVE algorithm. The observed scattering coefficient was well represented by the IMPROVE method reconstructed scattering coefficient had a regression coefficient of 0.90 and a correlation coefficient of 0.86 (Fig. 2c). Aerosol non-sphericity is relatively high during winter in Lanzhou, indicating the presence of mineral dust aerosols (Tian et al., 2015). A stepwise multiple linear regression model (Chan et al., 1999) rather than IMPROVE was chosen to reconstruct the aerosol absorption coefficient because FS and absorbing OC contributed to aerosol absorption in addition to EC in

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**Fig. 1.** Reconstruction of PM\(_{2.5}\) (a), comparisons between EC and BC at 880 nm, and relationships between ammonium and the sum of sulfate and nitrate (c).

**Fig. 2.** Linear relationship between the measured \( b_{\text{abs}} \), and EC (a), the measured \( b_{\text{abs}} \), \( b_{\text{scat}} \), \( b_{\text{ext}} \) and the reconstructed \( b_{\text{abs}} \) (b), \( b_{\text{scat}} \) (c), and \( b_{\text{ext}} \) (d).
Lanzhou. The regression equation was established as follows:

$$b_{abs} = 21.2982 \times [EC] + 1.4065 \times [FS] + 0.2123 \times [OC]$$  \hspace{1cm} (13)

The reconstructed absorption coefficient reasonably represented the observed absorption coefficient with a regression coefficient of 0.84 and a correlation coefficient of 0.94 (Fig. 2b). The regression coefficient and correlation coefficient between the reconstructed and observed extinction coefficient (i.e., the sum of scattering and absorption coefficient) were 0.91 and 0.89, respectively (Fig. 2d). Thus, the reconstruction method used in the present study is applicable in Lanzhou.

3. Results

3.1. Overview of aerosol optical properties during winter in Lanzhou

The aerosol optical parameters in the winter of 2019–2020 in Lanzhou were analyzed to better understand the aerosol optical properties in this region (Fig. 3). The range of the highest frequencies for $\sigma_{ap}$, $\sigma_{sp}$, and SSA were $15–30 \text{ Mm}^{-1}$, $350–400 \text{ Mm}^{-1}$, and $0.88–0.90$, respectively. Single scattering albedos $<0.7$ were correlated with samples with high $\sigma_{ap} > 200 \text{ Mm}^{-1}$. The average $\sigma_{ap}$, $\sigma_{sp}$ and SSA values were $64.70 \text{ Mm}^{-1}$, $363.68 \text{ Mm}^{-1}$, and 0.85, respectively (Table S2). Wintertime SSAs are relatively low in Lanzhou compared with eastern Chinese cities such as 0.94 for winter Hong Kong (Wang et al., 2017b), 0.89 for winter Nanjing (Zhuang et al., 2017), and 0.92 for hazy conditions during winter in Beijing (Zhang et al., 2020b).

The MAE$_{PM2.5}$ (MSE$_{PM2.5}$) ranged from 0.29 to 2.15 (0.55–8.78), with an average of 0.86 ± 0.31 (5.09 ± 0.70). Relatively high MAE further confirmed high absorption of atmospheric aerosols. AAE (SAE) ranged from 0.93 to 2.35 (0.31–3.89), with an average of 1.80 ± 0.18 (1.42 ± 0.28). The wide range of AAE and its average value of 1.80 indicated more other species in addition to BC aerosols. The aerosol hygroscopicity was very low (f(RH) = 1.68 ± 0.51), which might lead to different behavior of aerosol scattering coefficient as a function of PM against results in East China where aerosol scattering coefficient exhibits an exponentially growth with PM (Cheng et al., 2015; Ma et al., 2020). The $\Delta F/\tau$ was $22.58 \pm 3.13 \text{ W m}^{-2}$ in the winter of 2019–2020 in Lanzhou, the absolute value of which was less than that of $24.9 \text{ W m}^{-2}$ in Nanjing (relatively heavily polluted site; Shen et al., 2018) and higher than that of $19.9 \text{ W m}^{-2}$ in Finland (relatively clean site; Virkkula et al., 2011).

The scattering and absorption coefficients showed a clear double-
peaked diurnal variation, with the first peaks at noon (σ_{ap} at 11:00 and σ_{sp} at 13:00) and the second in the evening (20:00), and the first troughs before sunrise, with the second in the afternoon (Fig. 4a). The diurnal variation of PM_{2.5} chemical species in the winter of 2019–2020 in Lanzhou has also been studied recently (Du et al., 2020), which provides information needed to explain the diurnal variation in the optical parameters. The first troughs resulted from a lack of photochemically generated aerosols (Zhang et al., 2017; Du et al., 2020) and continuous diffusion during the night while the second were caused by favorable diffusion conditions (i.e., deep boundary layer height and high wind speed; Fig. S4). DAYTIME anthropogenic emissions, photochemical generations, and increasing boundary layer heights (Xu et al., 2016; Tan et al., 2017; Du et al., 2020) led to the first peaks, while vehicle emissions during the evening rush hour and unfavorable diffusion conditions (i.e., shallow boundary layer height and low wind speed; Fig. S4) facilitated the second ones. The scattering and absorption coefficients began to increase at 07:00 due to daytime anthropogenic emissions. Meanwhile, the boundary layer height rapidly increased after 08:00, leading to favorable diffusion conditions around noon. The accumulation of anthropogenic emissions and increasing diffusion conditions jointly resulted in a peak in the aerosol absorption coefficient at 11:00, when the concentration of EC also peaked (Du et al., 2020). However, the scattering coefficient continued to increase until 13:00, indicating a high concentration of photochemically generated scattering secondary inorganic aerosols (SIAs) around noon. Both the scattering and absorption coefficients peaked at 20:00, which further demonstrated the contribution of photochemical generations for the first peak in the scattering coefficient.

Single scattering albedo (MAE_{PM2.5}) exhibited low (high) values during the absorption coefficient troughs. Smaller particles from photochemical generation during the day led to lower MSE_{PM2.5} and higher SAE values during daytime. The diurnal variation of AAE was controlled by the aging of BC and changes in aerosol composition. Fresh BC in the late afternoon led to the lowest AAE at 19:00. Then aging of BC caused increasing trend in AAE until the next morning. Fresh BC emitted during the day led to a gradual decrease in AAE (see also Section 4.1). The diurnal variation of f(RH) was like that of RH (Fig. S1) but differed from that of the scattering coefficient, indicating the weak influence of humidity on aerosol scattering during winter in Lanzhou. The diurnal variation of ΔF/τ was consistent with that of the absorption coefficient, which implied that aerosol radiative forcing was strongly influenced by absorbing aerosols.

3.2. Aerosol scattering and absorption from low to high particulate pollution

The scattering and absorption coefficients generally increased with increasing PM_{2.5} in the winter of 2019–2020 in Lanzhou (Fig. 5a and b). The scattering coefficient exhibited a positive linear relationship with PM_{2.5}. However, the absorption coefficient showed two distinct linear relationships when PM_{2.5} was either lower or higher than 69 μg m\(^{-3}\). The absorption coefficient increased more rapidly with PM_{2.5} under high particulate pollution conditions. The SSA remained stagnant when PM_{2.5} was <69 μg m\(^{-3}\) but began to decrease rapidly with increases in PM_{2.5} under high particulate pollution conditions (Fig. 5c). The SSA for the highest PM_{2.5} bin reached as low as 0.82, which was lower than most cities in eastern China and globally under heavily polluted conditions such as 0.91 for Wuhan (Liao et al., 2020), 0.93 for Nanjing (Shen et al., 2018), 0.93 for Delhi (Tiwari et al., 2015), 0.91 for Finland, and 0.93 for Canada (Schmeisser et al., 2018). The MAE_{PM2.5} showed little variation for the lowest four PM_{2.5} bins but began to increase rapidly when PM_{2.5} was higher than 69 μg m\(^{-3}\) (Fig. 5f). Rapid increase of aerosol absorption coefficient and mass absorption efficiency when PM_{2.5} exceeded 69 μg m\(^{-3}\) seemed to be caused by the rapid increase of absorbing aerosols under high particulate pollution conditions (Fig. 6).

The scattering Ångström exponent exhibited little variation in the
lowest four PM$_{2.5}$ bins but began to increase when PM$_{2.5}$ exceeded 69 $\mu$g m$^{-3}$ (Fig. 5d), indicating increasing fine aerosols under high particulate pollution conditions. This phenomenon is opposite to that observed in eastern Chinese cities, where aerosols grow to larger size under polluted conditions due to aqueous reactions and hygroscopicity (Wang et al., 2016). In Lanzhou, the absorption Ångström exponent generally increased with PM$_{2.5}$ and began to decrease in the highest four PM$_{2.5}$ bins (Fig. 5e; see also Section 4.1).

The hygroscopicity of aerosols was investigated to better understand the aerosol scattering properties from low to high particulate pollution in the winter of 2019–2020 in Lanzhou. The correlation coefficient of the curve fitting relationship in Lanzhou City was as low as 0.65. The average RH was just 49.8% and RH rarely exceeded 80% during the winter of 2019–2020 in Lanzhou and such low humidity led to limited scattering enhancement of aerosols. The hygroscopic growth factor $f$(RH) decreased from 2.36 in the lowest PM$_{2.5}$ bin to 1.18 in the highest PM$_{2.5}$ bin, which was attributed to the relatively increasing hydrophobic OM and decreasing hydrophilic SIAs (Fig. 7b). In addition, the RH did not continue to increase with increases in particulate pollution (Fig. S3). Overall, the hygroscopic growth factor decreased with increases in PM$_{2.5}$, the weak influence of humidity on scattering led to an almost linear relationship between aerosol scattering coefficients and PM$_{2.5}$. This phenomenon is opposite to that recorded in eastern Chinese cities, where aerosol hygroscopicity and the generation of SIAs increase with increasing particulate pollution (Cheng et al., 2018; Wang et al., 2018; Ma et al., 2020) and SSA increases with increasing PM$_{2.5}$ (Zhang et al., 2020b).

3.3. The influence of aerosol composition on scattering and absorbing properties

The contribution of chemical species to aerosol extinction was studied using the reconstruction results (Fig. 8). The average reconstructed aerosol extinction coefficient was 475.0 Mm$^{-1}$ and OM was the
greatest contributor (34.0%), followed by ammonium nitrate (22.2%), ammonium sulfate (18.1%), EC (10.4%), and FS (3.7%). The contributions of OM, ammonium nitrate and EC increased from low (32.3%, 19.1%, and 9.4%) to high (35.7%, 22.7%, and 11.6%) pollution level, while the contribution of ammonium sulfate was the greatest for moderate level (20.3%).

The contributions of chemical species in Lanzhou to aerosol extinction were consistent with the results of some urban studies, such as in Beijing (Tao et al., 2015) and Shanghai (Han et al., 2015), where OM made the greatest contribution to extinction; the same conclusion was reached in Suzhou (Tian et al., 2016) for fine particles. Meanwhile the largest contributions of ammonium sulfate were found in Hainan (Tian et al., 2020), Guangzhou (Tao et al., 2014), and Xi’an (Cao et al., 2012).

The contributions of EC, FS, and OC on aerosol absorption were investigated in details (Fig. 8c and d). The average reconstructed aerosol absorption coefficient was 63.3 Mm\(^{-1}\) and the contributions of EC, FS, and OC were 77.4%, 16.6%, and 6.0%, respectively for total samples in the winter of 2019–2020 in Lanzhou. From low to high pollution levels, the contribution of EC absorption increased from 68.3% to 80.5%, while that of FS absorption decreased from 25.5% to 13.9%. The contribution of OC to aerosol absorption was minimal and exhibited little variation from low to high pollution levels.

4. Discussions

4.1. Combined effect of EC and FS on aerosol absorbing parameters

Single scattering albedo is a widely used parameter for aerosol absorption; the AAE is also an indicator of aerosol absorption. The AAE for fresh BC is ~1 (Bond and Bergstrom, 2006; Bond et al., 2013), while...
those for other absorbing particles and BC coated by non-absorbing materials are $>1$ due to varying absorption in different wavelength range (Lack and Cappa, 2010; Valenzuela et al., 2015; Devi et al., 2016). For example, the AAE for mineral dust aerosols exceeds 2.0 (Weinzierl et al., 2011) and that for BrC can reach as high as 9.5 (Lack and Langridge, 2013). The AAE in the present study varied from 0.91 to 2.92, with an average of 1.80 (Fig. 9). Overall, the SSA was the highest when the EC/PM$_{2.5}$ mass fraction was the lowest and AAE had the widest range of variations under this situation. With the increase of EC/PM$_{2.5}$, SSA decreased to below 0.80 and the range of AAE narrowed, converging to $\sim 1.65$.

Three parts of the SSA–AAE plots were selected to better understand aerosol optical properties during winter in Lanzhou: (A) the upper left with AAE $< 1.63$ and SSA $> 0.85$, (B) AAE $> 1.96$ and SSA $> 0.85$, and (C) SSA $< 0.80$. The sample numbers for parts A, B, and C were 194, 184, and 207, respectively. Part A of the SSA–AAE plots exhibited high SSAs and low AAEs. NH$_4$NO$_3$ accounted for the highest fraction (33.4%) of extinction for part A, followed by (NH$_4$)$_2$SO$_4$ (28.9%), and OM (21.1%). Both SSA and AAE were high in part B. The highest three extinction contributors were OM (38.7%), (NH$_4$)$_2$SO$_4$ (23.1%), and NH$_4$NO$_3$ (16.5%). Part C was characterized by low SSAs and medium AAEs. Organic matter contributed most to extinction (41.4%), followed by NH$_4$NO$_3$ (16.5%), and EC (15.6%). Relatively, part A was the cleanest and part C was the most polluted condition. There seems to be a gradual transition, first from A to B and then from B to C.

Elemental carbon and FS were the dominant species of aerosol absorption. The relatively low AAE and high SSA in part A were attributed to the low EC/PM$_{2.5}$ ratio. From part A to B, the contribution of FS (OM) increased more rapidly than that of EC, leading to an evident increase in the AAE and slight decrease in the SSA. From part B to C, the rapid increase in the contribution of EC, 8.4–15.6%, led to a large decrease in both the SSA and AAE. Since EC accounted for 77.4% of the aerosol absorption during the winter of 2019–2020 in Lanzhou, the SSA seems to be positively correlated with aerosol absorption, while the AAE was determined by the relative contribution of EC and non-EC absorbing species (i.e., FS and OM).

### 4.2. Influence of aerosol absorption on radiative forcing efficiency

The aerosol radiative forcing efficiency ($\Delta F/\tau$) determines the absolute value of radiative forcing. The relationships between SSA and parameter b of radiative forcing efficiency were investigated (Fig. 10), and it was found that $\Delta F/\tau$ strongly depended on SSA while no significant relationship was found between $\Delta F/\tau$ and b. This indicates that the aerosol cooling efficiency is mainly determined by SSA and, to a lesser extent, by particle size during winter in Lanzhou. This phenomenon is clearly different from the results in Nanjing, where $\Delta F/\tau$ is less correlated with SSA and more dependent on b (Shen et al., 2018). Aerosol absorption exerts a critical influence on the aerosol radiative forcing efficiency; this issue warrants further study.

### 4.3. Comparison of aerosol absorption among cities across the world

Aerosol absorption is important for aerosol radiative forcing. Two parameters, SSA and the absorption coefficient, were used to summarize the aerosol absorption in different regions of China and across the world in recent years (since 2015) (Fig. 11). The SSA was relatively high and the absorption coefficient was low for low PM levels during the winter of 2019–2020 in Lanzhou. However, the SSA decreased and the absorption coefficient increased from low to high PM level during the same period. The absorption coefficients of low and moderate PM levels were comparable to those of cities in other regions of China (Cheng et al., 2015; Deng et al., 2016; Wang et al., 2017a; Shen et al., 2018; Liao et al., 2020; Kong et al., 2020) and higher than those of European and American cities (Lim et al., 2018; Schmeisser et al., 2018; Donateo et al., 2020; Venter et al., 2020). The absorption coefficient at high PM level in Lanzhou was even higher than that of Delhi, Indian (Tiwari et al., 2015). Furthermore, the SSA in Lanzhou was much lower than that in Delhi, indicating strong aerosol absorption in Lanzhou. The sample number of the high PM level accounted half of the total samples.

Aerosol absorption coefficients are largely related to aerosol loadings, while SSAs are determined by the relative contributions of light scattering and absorption. In Sanya, a tourist city in China, both the absorption coefficient and the SSA are low (S1 in Fig. 11). The low

![Fig. 9. Relationships among optical parameters (SSA and AAE), chemical compositions, and pollution levels.](image-url)
The pollution level in Sanya results in a low absorption coefficient. The traffic emissions and biomass burning are the main sources of air pollution in Sanya in addition to the ocean emissions, which cause a relatively high contribution of EC (15%) and low contribution of SIAs (64%) to the aerosol extinction (Tian et al., 2020) compared to other cities. Thus, the SSA is also low in Sanya. In the suburbs of Hong Kong, the absorption coefficient is low and the SSA is high (P2 in Fig. 11). Similarly, the low absorption coefficient is related to low air pollution. However, the scattering aerosols (SIAs and OM) account for as much as 70.4% of the mass concentration of PM$_{2.5}$ in Hong Kong (Wang et al., 2018), which leads to strong scattering and relatively high SSAs. In Delhi, one of the most polluted cities in the world, both the absorption coefficient and SSA are high (F1 in Fig. 11). High air pollution leads to a high absorption coefficient and the average wintertime PM$_{2.5}$ concentration can reach 195.3 μg m$^{-3}$ (Tiwari et al., 2015). Secondary aerosols are the most important sources of pollution (Sharma et al., 2017) and scattering aerosols (SIAs and OM) account for ~75% of the mass concentration of PM$_{2.5}$ (Tobler et al., 2020), which leads to high SSAs in Delhi. The absorption coefficient is high and SSA is low in the present study of Lanzhou for the high pollution level (H in Fig. 11), which may be explained by the relatively high contribution of EC and low contribution of SIAs to PM$_{2.5}$ under high pollution during winter (Section 3.2 and Du et al., 2020).

5. Summary and conclusions

To clarify the evolution of aerosol absorption and quantify the contribution of absorbing species under different severities of particulate pollution, aerosol absorption from low to high particulate pollution was investigated via intensive observations of aerosol optical properties during the winter of 2019–2020 in Lanzhou, Northwest China.

The scattering coefficient linearly increased with increasing PM$_{2.5}$ while the absorption coefficient increased more rapidly when PM$_{2.5}$ exceeded 69 μg m$^{-3}$, leading to a rapid decrease in the SSA and sharp increase in MAE$_{PM_{2.5}}$ under high particulate pollution. Single scattering albedo (MAE$_{PM_{2.5}}$) decreased (increased) from 0.87 (0.76) in the lowest PM$_{2.5}$ bin to 0.82 (1.11) in the highest PM$_{2.5}$ bin. Low humidity led to limited aerosol scattering enhancement and the linear relationship between the scattering coefficient and PM$_{2.5}$ was attributed to decreases in aerosol hygroscopicity with increasing PM$_{2.5}$.

Elemental carbon, FS, and OC accounted for 77.4%, 16.6%, and 6.0% of the total aerosol absorption, respectively. From low to high pollution levels, the contribution of EC absorption increased from 68.3% to 80.5% while that of FS decreased from 25.5% to 13.9%.

The calculated aerosol radiative forcing efficiency was strongly correlated with SSA, showing the importance of aerosol absorption for aerosol radiative forcing. Our results show the unique behavior of a rapid increase in aerosol absorption under high particulate pollution during winter in Lanzhou, which is opposite to the trends observed in

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Fig. 11. Comparison of aerosol absorption in Lanzhou and other sites.
eastern Chinese cities, where SSA increases with increasing PM$_{2.5}$. However, the radiative forcing of light-absorbing materials (i.e., BC, mineral dust, and BrC) must be quantified in future studies.

CRediT authorship contribution statement


Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgement and data availability

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Appendix A. Supplementary data

Supplementary data to this article can be found at https://doi.org/10.1016/j.atmosenv.2020.118081.

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